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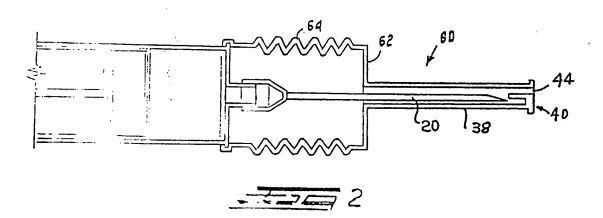
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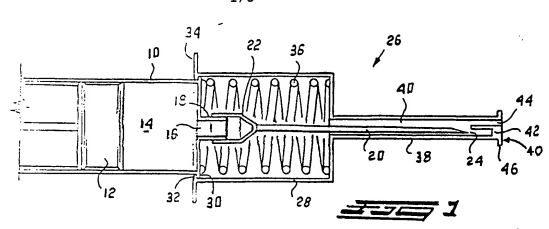
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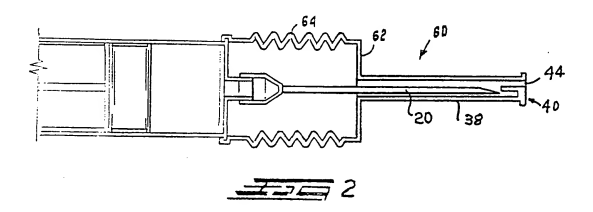
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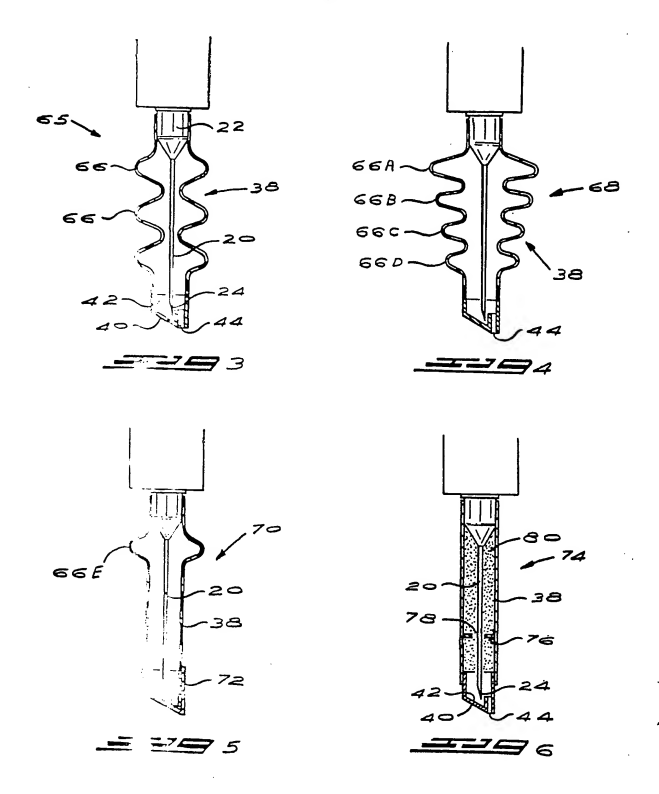
(54) Syringe needle or catheter covers

(57) A protection in the 60 for a needle 20 of a syringe or catheter has a member 38 within which the needle point can be located. A blocking surface 40 at one end of the member opposes the needle point and has a hole 44 which can be brought into register with the point to allow the needle to extend from the member. The member may be tubular and secured to the syringe by a socket portion 62 formed with resilient bellows 64 or be of a more rigid tubular construction and mounted for telescopic monoment over the syringe against the action of a coil spring (Figure 1). In further embodiments (Figures 3-5) the tubular mile hards mounted on the needle hub and comprises at least one resilient bellows formation. In another end admirant, and a) a resiliently flexible tubular member is filled with low density foam rubber for enhanced resilience. In yet another either the figure 7) a needle tip protecting cap is attached to the needle hub by a resilient strip.

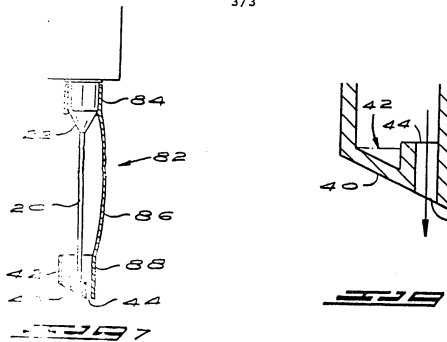


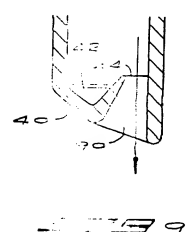


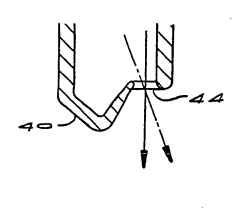




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This invention is concerned with a protective device for a needle of a syringe, catheter or similar medical appliance for human or animal use.

The invention provides a protective device for an elongate needle of the kind described which has a base at one end and a point at an opposing end, the device including a blocking surface and means which extends from the base and which supports the blocking surface, the blocking surface having a hole and being movable at least towards the base to allow the needle to extend through the hole.

The device may be provided so that the needle is aligned with the hole, but not extending therethrough. The device is then ready for use. After use the blocking surface is brought automatically or manually to a position at which it opposes the needle point.

Alternatively the device is provided with the blocking surface opposing the new point and the blocking surface is movable, preferably laterally, relatively to the needle point to bring the hole and the needle point into allegement, and thereby ready the device for use.

The support means may take on any suitable form and may comprise one or the strips of material, which may be resiliently deformable, and which

extend from the base to the blocking surface.

The support means may alternatively include a tubular member within which the needle is located with the blocking surface at one end of the tubular member.

A disc may be located inside the tubular member and may have an aperture through which the needle extends to maintain the needle point out of register with the said hole.

The tubular member may be made from a resiliently deformable and medically acceptable material such as a rubber or plastics material, foam rubber or the like, and preferably is transparent. A deformable filler material such as sponge rubber may be inside the tubular member. Sealing means may be engaged with the tubular member to keep the need to sterile. The sealing means may be a dust cap, or a stopper which fits into the hole, or the like.

The tubular member may include at least one deformable bellows formation which may be near the base of the needle, or may include a plantary of spaced bellows formations of the same or different sizes.

The device may include biasing means which acts against the support

means when the blocking surface is moved towards the base and thereafter acts to extend the support means automatically to its initial position.

The blocking surface may include a recessed formation in the form of a blind or dead-end passage which may oppose or which may be moved so that it opposes the point of the needle and the hole may be formed adjacent the recessed formation. Movement of the needle towards the blocking surface thus causes the point to enter the dead-end passage. The blocking surface is of a material which is sufficiently hard or thick to prevent penetration of the blocking surface by the point. The blocking surface may include surfaces which diverge away from the hole so that which the needle extends through the hole it is completely free of the blocking surface. The recessed formation may surround the hole.

The support means preferably engages directly with the base or hub of the modele but may engage with a syringe or other instrument with which the modele is used.

The blocking surface may be moved laterally and twisted relatively to the mounte to pring the hole into register with the point or, conversely, to move the hole so that it is not aligned with the point.

The invention is further described by way of examples with reference to the accompanying drawings in which:

Figures 1 to 7 are side views in cross section illustrating different embediments of the protective device of the invention, and

Figures 3 to 10 are side views in cross section respectively illustrating in enlarged detail different possible forms of construction which can be embinated in the protective device.

Figure 1 illustrates one end of a known syringe 10 which includes a plant of 12 which is adapted to expel fluid 14 of any appropriate kind through a nozzle 16 formed by a spigot 18. A stainless steel injection not 1.00 with a base or hub 22 is frictionally engaged in a leakproof manner with the spigot 18 so that the contents of the syringe are, on activation of the plunger, expelled through a leading end or point 24 of the need.

A principle device 26 is engaged with the syringe and comprises a housing formed from a clear plastics material which is slightly resiliently decomposed. The housing has a socket 28 with a flange 30 which engages with a undercut formation 32 on the socket.

The socket 28 includes two opposed extensions 34 or, alternatively, a continuous flange 34. If manual force is exerted on the formation 34 then the socket can be moved to the left relatively to the syringe. A coil spring 36 inside the socket acts between a right hand end of the socket and an opposing end of the syringe.

The accusing includes a tubular member 38 within which the needle 20 is 10.000 d. At one end the member has a blocking surface 40 with a blind or double and passage 42 and a hole 44 adjacent the passage 42. Under normal conditions the needle 20 is aligned with the dead-end passage with the point 24 at the mouth of the passage and if the device is moved to 10 left relatively to the syringe, the point 24 is advanced into the 10.000

the fible 44 is aligned with the needle point. As an injection is given the needle point penetrates the skin of the recipient. A reaction force is greated on the outer side of the blocking surface and this causes the handing to move to the left relatively to the syringe, against the bias of the spin grade.

the needle is withdrawn the tubular member automatically moves
to right under the action of the spring. The emerging needle is

there is a fully and automatically enclosed by the tubular member and the nature silience of the tubular member re-aligns the needle point with the continend passage. This minimizes the likelihood of any infection being a sidentally transferred via the needle.

Figure 1 shows a device with a housing 60 which has a socket 62 formed with 1 10 ws 64 which have a natural resilience resulting from their shape and 1 1 wial. When an injection is given and the member 38 is moved to the 10 the bellows are compressed. When the needle is removed from their cibient the bellows automatically expand and the tubular member 38 is 10 1 d to the right to enclose and protect the needle point.

Fig. 3 illustrates a simplified device 65 which comprises a tubular may 1 38 made from a resilient material such as latex rubber with a plu 1 of spaced bellows formations 66. At one end the tubular member end; 3 directly with a hub 22 of an injection or catheter needle 20. The policy is of the needle lies at the mouth of a dead-end passage 42 on a bid 1 surface 40 at an opposing end of the tubular member. A hole 44 is 1. It adjacent the passage.

the letter away from the hub and is displaced laterally to align the product the hole 44. As the needle enters the recipient the member

the needle is withdrawn the formations 66 automatically extend the many or 33 and the needle is thereby retracted into the member 38 and the needle is the passage 42.

Fig. 4 shows a device 68 which is similar to the device 65 but with the similar to the similar to the device 65 but with the similar to the similar to the similar to the device 65 but with the similar to the

5 shows a device 70 with a single bellows formation 66E and with malacter of the tubular member 38 being of more or less constant section. A removable dust cap 72 encloses the outer end of the arction that the needle 20 is kept in a sterile condition within the ar. This tubular member is easier to make than the members of \$3 ard 4. Instead of the dust cap a stopper can be plugged into the arctic seal the interior of the tubular member.

another in any appropriate way, namely the member 38, and the secured suggestation which is made from a relatively hard plastics material that leedle point does not readily penetrate. It is preferred to the tubular member with the base of the needle but it is possible to the member to a syringe or other device with which the needle is

used.

Figure 3 shows a device 74 with a tubular member 38 of regular cross section made from a resiliently flexible material such as foam rubber. A disc 73 is positioned inside the member and has an aperture 78 through which the needle 20 passes. The disc keeps the point 24 aligned with a dead-and passage 42 on the blocking surface 40. The member 38 can be defied to one side so that the hole 44 is aligned with the point 24 when an including is to be given.

This is after of the member 38 may optionally be filled with an easily comparable material 80, such as low density foam rubber, to enhance its assume. Alternatively the resilience can be achieved from the foam rubb. It also be and the foam rubber can be protected by means of the member and which can be made from an impermeable material.

A discussion of the disc 76, can also be used with the embodiments of Fig. : 3×5 .

Figure 7 shows a device 82 which is made as a one-piece plastics mouth 1g. A ring 84 is engaged with the hub 22 of a needle 20 and at least call to 21 material 86 extends from the ring to a member 88 which call a dead-end passage 42 and a blocking surface 40 with a hole

41 hount the passage 42.

The trip 85 aligns the passage 42 with the point 24. When an injection is to be given the strip is deflected laterally so that the point 24 is aligned with the hole 44. The member 88 is movable along the needle with the strip 35 deflecting outwardly away from the needle as an injection is given. Once the needle is withdrawn the strip 86 returns to its original and contained its own resilience and the needle point is withdrawn the passage 42.

The point is automatically protected after an injection is given. This was accessarily be the case for the protective device can be made a maderial which has a low natural resilience. The onus then rests on that per to cause it to extend so that the needle is brought to a protected process with its point opposing the blocking surface, after an injection that peen given.

and 10 illustrate in enlarged cross section different forms of must on for the blocking surface 40. In Figure 8 the blocking surface to the axis of the needle and the hole 44 adjacent the dead-end defined by a short passage 90 of circular cross section. In

44 c. Ins into a passage 90 of increasing cross section. In Figure 10 on the error hand the hole 44 does not lead to a passage 90 of the kind shown in Figures 8 and 9. The two latter constructions prevent the point of the needle from inadvertently snagging on the blocking surface.

With the devices of Figures 1 to 7 the point of the needle is not initially aligned with the hole, but rather with the dead-end passage. It is possible, passed larly with the embodiments of Figures 3 to 5, to provide the device with the needle point aligned with the hole 44, partly inside the passage 90, and ready for use. After an injection is given the blocking surface can be a read canually so that the needle point opposes the dead-end passed and is then protected. It has also been found that when the resilience of the member then causes movement of the blocking surface so that the point is fully retracted from the passage 90 and the resilience of the member then causes movement of the blocking surface so that the point is aligned with the dead-end passage, and is protected. As an effect is achieved simply by shaking the syringe - the movement can be dead-end passage.

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- A protective device for an elongate needle with a base at one end and a point at an opposing end, the device including a blocking surface and means which extends from the base and supports the ling surface, the blocking surface having a hole and being movable at a st towards the base to allow the needle to extend through the hole.
- A protective device according to claim 1 wherein the support sometimes at least one strip of material which extends from the to the blocking surface.
- A protective device according to claim 1 wherein the support us includes a tubular member within which the needle is located and locking surface is at one end of the tubular member.
- A protective device according to claim 3 which includes a cis which is located inside the tubular member and which is formed with persure through which the needle extends.
- A protective device according to claim 3 or 4 wherein the

- 6. A protective device according to claim 3, 4 or 5 wherein the tubular member includes at least one deformable bellows formation.
- 7. A protective device according to claim 1 which includes biasing means which acts against the support means when the blocking surface is moved towards the base.
- 8. A protective device according to any one of claims 1 to 7 wherein the blocking surface includes a recessed formation which oppose a the point and the hole is formed adjacent the recessed formation.
- 9. A protective device according to any one of claims 1 to 8 which is provided with the needle point aligned with the hole.
- A protective device according to any one of claims 1 to 8 which is provided with the needle point opposing the blocking surface.
- 11. A protective device substantially as hereinbefore described with reserve to any one of the accompanying drawings.

Patents Act 1377

F taminer's report to the Comptroller under Section 17 (The Search Report)

Application number

Relevant Tech: pal fields	Search Examiner
(i) UK CI (Editio . K) ASR (RGG), (RGM)	
(ii) Int CI (Editio: 5) A61M	R J WALKER
Databases week werk	Date of Search
(i) UK Patent C to be	
(ii)	10 APRIL 1992
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Documents considered relevant following a search in respect of claims

Category (see ove:	Edentity of document and relevant passages		
X,P	:3 2243552 A	(TRANSFERTEC LTD) see eg page 13 lines 14-25	1,3, 7-10
X,P	12 0434008 A	(LAZOVSKI) see Figures 1 and 2	1,2,7,10
X,P	13 0413 872 A	(BOISSON-MULLER) see drawings	1,3-10
x	0 80 10767 A	(DEEKS) see eg Figures 7 and 8	1,2, 7-10
X,P	5015 240	(SOOPRONI ET AL) see abstract	1,3,5,6, 7,9,10
x	US 4978344	(DOMBROWSKI ET AL) see Figures 5-7	1,3,6,8,
x	1 4915366	(COREY) see Figures 8 and 9	1,2,3,4,9,10
x	11.40 0521	(TAICO ET AL) see whole document	1-6,9
x	3 4725267	(VAILLANCOURT) see whole document	1-4, 6-10

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Category	Identity of document and relevant passages	Relevant to claim(s
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